# This Page Is Inserted by IFW Operations and is not a part of the Official Record

# **BEST AVAILABLE IMAGES**

Defective images within this document are accurate representations of the original documents submitted by the applicant.

Defects in the images may include (but are not limited to):

- BLACK BORDERS
- TEXT CUT OFF AT TOP, BOTTOM OR SIDES
- FADED TEXT
- ILLEGIBLE TEXT
- SKEWED/SLANTED IMAGES
- COLORED PHOTOS
- BLACK OR VERY BLACK AND WHITE DARK PHOTOS
- GRAY SCALE DOCUMENTS

# IMAGES ARE BEST AVAILABLE COPY.

As rescanning documents will not correct images, please do not report the images to the Image Problem Mailbox.

## PATENT SPECIFICATION

1 585 661 (11)

(21) Application No. 23718/76

(22) Filed 8 June 1976

(21) Application No. 23734/76

(22) Filed 8 June 1976

(21) Application No. 23736/76

(22) Filed 8 June 1976

(23) Complete Specification filed 9 June 1977

(44) Complete Specification published 11 March 1981

(51) INT CL3 C07D 498/04 A01N 43/90 A61K 31/42 (C07D 498/04 205/00 263/00)

(52) Index at acceptance

C2C 1315 214 220 226 22Y 247 255 25Y 292 29Y 306 30Y 351 352 360 361 366 367 368 36Y 387 388 490 491 623 625 628 652 658 65X 678 699 761 762 801 805 80Y AA TR

(72) Inventors EUNICE JEAN NAPIER JAMES REGINALD EVANS DAVID NOBLE MICHAEL EDWARD BUSHELL GRAHAM WEBB and

DAVID BROWN

SCIENCE REFERENCE LIBRARY

5

10

15

20

25

(54) CLAVAM DERIVATIVES

(71)We, GLAXO LABORATORIES LIMITED, a British Company of Greenford, Middlesex, England, do hereby declare the invention, for which we pray that a patent may be granted to us, and the method by which it is to be performed, to be particularly described in and by the following statement:—

This invention relates to novel antibiotic compounds and to a process for their

production.

5

10

15

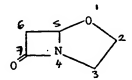
25

Fermentation of Streptomyces clavuligerus, and in particular strain NRRL 3585, is known to produce a number of antibiotic substances and British Patent Specification No. 1,315,177 describes and claims the cultivation of Streptomyces clavuligerus strain NRRL 3585 until a substantial amount of two β-lactam carboxylic acids, referred to as Antibiotics A 16886 I and A 16886 II is produced. In German OLS 2,604,697 we have described the isolation from such fermentation broths of a further B-lactam carboxylic acid, clavulanic acid.

We have now been able to isolate from fermentations of strains of Sterptomyces clavuligerus further  $\beta$ -lactam compounds which have been found to possess

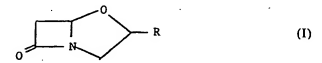
antibiotic activity.

The compounds of this specification are named with reference to "clavam", the name given to the parent heterocycle of formula A



by analogy with the term "cepham" used in the naming of cephalosporin compounds in J.Amer.Chem.Soc., 1962, 84, 3400. 20

According to one aspect of this invention, therefore we provide a compound of the formula (I)



wherein R represents a hydroxymethyl group, said compound in deuteroacetone exhibiting the 100 MHz proton n.m.r.  $\tau$  values shown in Table 1 herein, and the corresponding compounds in which R is a formyloxymethyl group or a carboxyl or esterified carboxyl group and, in the case where R represents a carboxyl group, salts thereof.

### TABLE I

	TA	BLE I	-
5	τ value	τvalue	5
-	4.68 (d, 2.5 Hz) (1H) 5.58 (multiplet) (1H)	7.09 (dd, 6 Hz and 11.5 Hz) (1H)	5=
	5.77 (broad singlet,	6.32 (broad singlet) (2H)	*
	exchanges with D <sub>2</sub> O) (1H)	672 (44.2 5115 and 1611 ) (11)	.2
	6.14 (dd, 7 Hz and 11.5 Hz) (1H)	6.72 (dd, 2.5 Hz and 16 Hz) (1H)	
	0.14 (dd, / f12 alid 11.5 f12) (1H)	7.29 (d, 16 Hz) (1H)	
10	It is believed that the - values of	direction Table 1	1000
	hereinaster are subject to an experime	given in Table 1 and in Tables 2 and 3	10
	It should be noted that although	the stereochemical configuration of the	
	compounds of the invention is not ke	the stereochemical configuration of the nown, the stated $\tau$ values in Table I are	
•	characteristic of the particular configur	alion existing in the commound of form 1.	
15	(1) III WILLIAM IS IIVUIOXVII A	nd all the compounds have the same	1.5
	comiguration at the 2 and 3 positions.		15
	Compounds of formula (1) in which	R is hydroxymethyl, formyloxymethyl or	
	esternied carboxyr snow userur and anti-	ilingal activity: the compounds inh. b	
20	is callowly and sails incled are primar	IIV Of IISE ID Dreparing active estare	
20	ine saits according to the invention	Include salts with inorganic bases, such as	20
	aikan metai sans, e.g. soqium, porassin	m and lithium calter alkaling ageth	
	saits, e.g. calcium and magnesium saits:	and ammonium salts as well as salts with	
	organic bases, for example amine saits.		
25	The esters according to the invent	ion may be represented as compounds of	
	which is conveniently a group —COC	OR' where R' represented as compounds of OR' where R' represents an organic group	25
			23
30 ·			
	the preparation of further esters.	ed forms of the parent acid, for example in	30
	Thus, the group R <sup>1</sup> may represent	a straight or branched unsubstituted or	
3.6	example, a methyl, ethyl, propyl or iso	propyl, butyl, sec-butyl tert-butyl or allyl	
35	group, desirable substituents being, for e	propyl, butyl, sec-butyl tert-butyl or allyl xample, alkoxy, e.g. methoxy; halogen, i.e.	
	fluorine, chlorine, bromine or iodine; of	syano; acyloxy, e.g. methoxy; halogen, i.e.	35
	acetoxy or pivaloyloxy, or alkoxycarbo	ryalio, acyloxy, e.g. alkanoyloxy, such as nyloxy, such as ethoxycarbonyloxy; acyl,	
40			
			40
	substitued, e.g. by a C <sub>1-4</sub> alkyl group, p		•
45	an aryl group having up to 12 carbon	atoms a glamband and but the	
			45
	a cylcoalkyl group containing not me	ore than 12 carbon atoms, e.g. adamantyl;	
50			
50	a heterocyclic group containing not	more than 12 carbon atoms, the hetero	60
	atom being, for example, oxygen, as in the	e tetrahydropyranyl or phthalidyl group	50
•	rotation [M123 in 4] in which	e tetranydropyranyl or phthalidyl group.  R is —CH <sub>2</sub> OH has a negative molecular	
	rotation [M] <sub>6</sub> <sup>23</sup> in dimethyl sulphoxide, n	amely -166°. The 100 MHz proton nmr	
55 .	spectrum of a solution of this compound shown in Table I, the full spectrum being	in deuteroacetone revealed \u03c4 values as	
- •	shown in Table I, the full spectrum beir drawings.	g snown in Fig. 1 of the accompanying	55
	The compound of formula (I) where	n D romania C	
	shows in deuterochloroform solution τ val as shown in Table ?	n R represents a formyloxymethyl group	
	as shown in Table 2.	des in the 100 MHz proton nmr spectrum	

these formulations, the concentration of active material is preferably between 0.1% and 40% by weight. The antifungal compounds of the invention may also be of use as storage preservatives in certain materials, for example, food, wallpaper paste, paint or 5 petrol, or in beer and wine to prevent undesirable fermentation. In addition, the compounds may be of use as seed dressings. The compounds of the invention may be isolated from a fermentation broth prepared by culture of a strain of Streptomyces clavuligerus. Particularly useful strains are Streptomyces clavuligerus strain NRRL 3585 and 10 mutants thereof. We have found strains NCIB 11260 and NCIB 11261 to be 10 especially useful. Strain NCIB 11260 is a single colony isolate from strain NRRL 3585 having essentially similar morphology to NRRL 3585, as described in British Patent Specification No. 1,315,177. Strain NCIB 11261 also has essentially similar morphology to strain NRRL 3585, except that it requires uracil for growth. 15 As used herein, the term 'mutant' will include any mutant strain which arises 15 either spontaneously or as a result of the action of an external agent, which may be either deliberately applied or otherwise. Mutant strains may be produced by a variety of methods including ionising radiation, chemical treatment and genetic techniques, such as those outlined in Techniques for the Development of Micro-20 Organisms by H. I. Adler in "Radiation and Radio-isotopes for Industrial 20 Microorganisms", Proceedings of the Symposium, Vienna, 1973, p. 241, International Atomic Energy Authority. In the preparation of NCIB 11261, we used  $\gamma$ -radiation, e.g. of about 80 kilorads, NCIB 11261 has been found to show a requirement of uracil for growth, 25 and the yield of the compounds of the invention has been found to be dependent to 25 some extent on the amount of uracil present in the fermentation medium. It is preferred that the level of uracil is not greater than 200  $\mu$ g/ml of broth, and preferably from 5 to 125  $\mu$ g/ml. The production of a compound of formula (I) wherein R is hydroxymethyl, 30 formyloxymethyl or carboxyl or a salt of a compound in which R is carboxyl by 30 fermentation of Streptomyces clavuligerus may be effected by conventional means, i.e. by culturing the Streptomyces clavuligerus in the presence of assimilable sources of carbon, nitrogen and mineral salts. Where a compound of formula I in which R represents an esterified carboxyl group is desired the corresponding compound in which R is carboxyl may be esterified either after isolation of the clavam carboxylic 35 35 acid or in situ followed by isolation of the desired ester. Cultivation will preferably be carried out by submerged culture under aerobic conditions. Assimilable sources of carbon, nitrogen and minerals may be provided by either simple or complex nutrients. Sources of carbon will generally include 40 glucose, starch glycerol, molasses, dextrin, lactose, sucrose, carboxylic acids, 40 alcohols, for example, methanol, n-paraffins and vegetable oils. Sources of nitrogen will generally include soyabean meal, corn steep liquors, distillers solubles, yeast extracts, cottonseed meal, peptones, casein, amino acid mixtures, ammonia (gas or solution), ammonium salts or nitrates. Urea and other 45 amides may also be used. 45 Nutrient mineral salts which may be incorporated into the culture medium include the generally used salts capable of yielding sodium, potassium, ammonium, iron, magnesium, zinc, nickel, cobalt, manganese, vanadium, chromium, calcium, phosphate, sulphate, chloride and carbonate ions. 50 An antifoam may be present to control excessive foaming and added at 50 intervals as required. Cultivation of the Streptomyces clavuligerus will generally be effected at a temperature of from 20°-32°C preferably of from 25-30°C, and will desirably take place with aeration and agitation, e.g. by shaking or stirring. The growth 55 55 medium may initially be inoculated with a small quantity of sporulated suspension of the microorganism but in order to avoid a growth lag a vegetative inoculum of the organism may be prepared by inoculating a small quantity of culture medium with the spore form of the organism, and the vegitative inoculum obtained may be transferred to the fermentation medium, or, more preferaby to one or more seed 60 stages where further growth takes place before transfer to the principal 60 fermentation medium. In a preferred embodiment of the fermentation a slope of Streptomyces clavuligerus may be used to inoculate a medium comprising sources of assimilable carbon, e.g. sucrose and/or glycerol, assimilable nitrogen, e.g. tryptones and/or complex mixtures of assimilable carbon and nitrogen, e.g. distiller's solubles and/or 65 65

The second second second second second

nective particular of the plant was sufficiently and the plant of the property of the plant of t

6	1,585,661	6 .	***
	required, to further fractionation. As described in detail hereinafter, the acid of the invention or a salt thereof may be subsequently eluted from the resin.  The anion exchange resin will generally carry amino groups (weakly basic) or quaternary ammonium groups (strongly basic). The resin may, for example, be a polystyrene polyscrylic enorgy polystyrene.		TAN BURNE
	dextran resin and may be macroreticular or microreticular. The term 'resin' is used herein for convenience also to include cellulosic derivatives and the above dextran derivatives which are derived from naturally occurred to be above dextran	5.	THE STREET, ST
10	tertiary amino groups), and Amberlite IRA 93 (Macroreticular: polyacrylate: linked with divinylbenzene: tertiary amino groups) both sold by Rohm & Haas (U.K.) Ltd. of Croydon, England. (Amberlite is a registered Trade Mark). Typical strongly basic ion exchange resins include Zerolit EF. Zerolit EF.	10	4、16.15(21.68)。18.14)。18.14)。18.14)
15	(Zerolit is a registered Trade Mark).  In another embodiment, the clarified or unclarified broth or other solution containing at least the hydroxymethyl and/or formyloxymethyl compounds of the invention may be applied to a non-ionic recip of the solution of the containing at least the hydroxymethyl and/or formyloxymethyl compounds of the	15	
20	pore/ml bead; sold by Rohm & Haas (U.K.) Ltd.) which does not retain the hydroxymethyl and formyloxymethyl compounds of the invention but which will retain several other significant broth components.  In a still further alternative, the hydroxymethyl and (a.g., a.g., a.	20	
25	an ester solvent such as ethyl acetate or an alcohol such as butanol. Such extraction may be applied to the clarified or unclarified broth, or to the eluates or effluents from the foregoing adsorption elution procedures if necessarily and the solution of the solution procedures if necessarily and the solution of the	25	
30	in the pH range 5—8, the acid according to the invention and similar acids, where present, will be in salt form and will remain in the aqueous phase. As indicated hereinafter, solvent treatment at acid pH enables the acid of the invention to be extracted.	30	
35	To permit extraction into a suitably small volume of water-immiscible solvent it may be desirable to concentrate the solution, e.g. by evaporation under reduced pressure. A high concentration of a salt such as ammonium sulphate assists the extraction.  The desired compound of formula (I) wherein R represents a hydroxymethyl or formyloxymethyl group may be further applicable.	35	
40	or an organic solvent-compatible, cross-linked dextran such as Sephadex LH 20 (sold by Pharmacia U.K. Ltd.). (Sephadex is a registered Trade Mark). The solution of the compound obtained from the previous purification stage may be too dilute for application to the column and may conveniently be concentrated by evaporation under reduced pressure.	40	
45	The column carrying the desired compound may then be eluted, for example using a solvent of suitable polarity. In the case of silica columns, ethyl acetate containing a hydrocarbon, e.g. hexane or toluene, may be used to elute the formyloxymethyl compound and the same solvent may be used to elute the hydroxymethyl compound.	45	
50	predominantly the compound of formula I in which R is a formyloxymethyl group and the compound of formula I in which R is a hydroxymethyl group is eluted in the later fractions.	. 50	
55	Finally, the fractions containing the desired compound may be combined and evaporated to yield the desired compound.  By a suitable combination of the foregoing procedures, the compounds in which R is a formyloxymethyl or hydroxymethyl group have been isolated as pale yellow oils of at least 90% purity. However, in this form the compounds are unstable and are best stored in solution in water or organic solvents.	55	•
60	As indicated above, the acid of the invention can be separated from the hydroxymethyl and formyloxymethyl compounds and other broth components by application of the clarified or unclarified broth or other aqueous solution containing the acid and/or a salt thereof to an animon exchange resin. Such a resin is	60	
65	conveniently in the salt form, e.g. the chloride form. The acid of the invention, usually together with one or more other $\beta$ -lactam carboxylic acids, notably clavulanic acid, may then be eluted from the ion-exchange resin, conveniently	65	_ _

	麦			
•	6	7	1,585,661	7_
of the	6 5		using an aqueous solution of a salt as eluant. We have found that solutions of lithium salts, e.g. at a molarity 0.2 to 2.5M, are particularly useful as eluants since lithium salts, e.g. at a molarity of a contaminating acids, clayulanic acid, can be	
sic) or . be a			the lithium salt of one of the principa to logge the lithium salt of the acid of formula	5
inked s used	5	5	I in the mother liquor. In general, the cluate containing salt of about 5—10,	
eakly	₩.		before precipitation of the lithium clavulation of till further lithium salt. Alternatively,	
ylate:	<u> </u>		the acid of the invention has been with an aqueen	10
cross- Haas	10: 🛁	10	However, in order to minimise elution of advolved impartition from the distance of the distanc	
pical	<u> </u>		high concentration. Alternatively, after elited impurities and the precipitate	
rnia).	15	15	separated off before further treatment. The solvent hand, isopropanol or ethylene	15
ution of the			glycol, an ether such as dioxan or tetranyuroida or dimethylsulphoxide. In general,	
AD-2 15 ml	}		alcohols are preferred as such solvents, e.g. entation of the alcohol thereto	20
1 the 1 will	20	20	being from 70 to 97% by volume. Oldeshable indexing the separated, e.g. by centrifugation or filtration. Fractional crystallisation as be separated, e.g. by centrifugation of filtration component of the supernatant, or	
ethyl , e.g.			described above, to remove all undesired major compension or a salt thereof filtrate, yields a mother liquor from which the acid of the invention or a salt thereof	25
ction ients	25	25	may be isolated.	
f any cted			exchange resin, the salts therein tend to overload the resin. It is stated a salts therein tend to overload the resin. It is stated as a salts therein tend to overload the resin. It is stated as a salts therein tend to overload the resin. It is stated as a salts therein tend to overload the resin. It is stated as a salts therein tend to overload the resin. It is stated as a salts therein tend to overload the resin. It is stated as a salts the resin tend to overload the resin. It is stated as a salts therein tend to overload the resin. It is stated as a salts therein tend to overload the resin. It is stated as a salts therein tend to overload the resin. It is stated as a salts therein tend to overload the resin. It is stated as a salts therein tend to overload the resin. It is stated as a salts the resin tend to overload	
here ated	20	30	adsorbent carbon, followed by entition with an adversariative thereof may be isolated	30
o be	30	, 30	from the mother liquor remaining after removal of the depend on whether or convenient means, but the method employed will, of course, depend on whether or convenient means, but the method employed will, of course, depend on whether or convenient means, but the method employed will, of course, depend on whether or convenient means, but the method employed will, of course, depend on whether or convenient means, but the method employed will, of course, depend on whether or convenient means, but the method employed will, of course, depend on whether or convenient means, but the method employed will, of course, depend on whether or convenient means, but the method employed will, of course, depend on whether or convenient means, but the method employed will, of course, depend on whether or convenient means, but the method employed will, of course, depend on whether or convenient means, but the method employed will, of course, depend on whether or convenient means, but the method employed will, of course, depend on whether or convenient means, but the method employed will, or course, depend on whether or convenient means are convenient means.	
vent	į		not it is the acid of the invention of a salt of ester the form substantially to free the Further purification means will be desirable in order substantially to free the compound of the invention from minor amounts of impurities which may have	35
the	35	35	been carried through the earlier purification steps.	
thyl ilica		}	liquor remaining after removal of the clavellande of solvent, for example	40
I 20 The	40	40	an ester such as etnyl acetate, a ketone such as methy	40
too l by	, , ,		alcohol such as butanol.  Alternatively, ion-pair extraction can be used, for example by extracting the aqueous medium containing the desired acid and/or salt thereof with a solution of aqueous medium containing the column (e.g., a hydrogarbon such as hexane or	
iple tate	45	45	an amine in a water-immiscible solvent (e.g. a hydrogenetal as methylene chloride). The amines kerosene or a halogenated hydrogenetal as methylene chloride). The amines	45
the one (20,			aliphatic groups, e.g. a branched chain plantaly of scottestite XLA3, LA1 and LA2 molecular weight range 280—400g for example, Amberlite XLA3, LA1 and LA2 molecular weight range 280—400g for example, Amberlite XLA3, LA1 and LA2	
ain oup	50	50	and Primene JMT (Sold by Rohm & Haas (O.R. Etd.), (Thinking and Primene JMT (Sold by Rohm & Haas (O.R. Etd.), (Thinking a 3—7) so that Trade Mark). The pH of the system should be acidic (e.g. in the range 3—7) so that Trade Mark). The pH of the system and the acid of the invention in the acid form. The	50
l in			Trade Mark). The pH of the system should be actide (e.g. in the late of the the amine is in the salt form and the acid of the invention in the acid form. The desired acid is extracted into the solvent phase to form a salt with the amine. The acid pH of the system may be achieved by addition of a mineral acid such as acid pH of the system may be achieved by addition of a mineral acid such as	
ınd		· 55	hydrochloric acid or an organic acid such as a second the organic solvent solution	55
in ale	55	\ .	into an aqueous medium containing an equeous salt solution.	
ire		ļ	Where an ester is desired, an extract of the for example diazomethane	<i>2</i> 0
.he by	60 .	. 60	or diphenyldiazomethane. The resulting solution has be too dilute for application	60
on : is	-		compound of the invention in an organic solvent, may be too didaction is to a column for purification purposes and will thus, if this mode of purification is desired, preferably be evaporated to dryness followed by redissolution into a desired, preferably be evaporated to dryness followed by redissolution into a	
)D		i ·	desired, preferably be evaporated to drylless followed by	

desired, preferably be evaporated to dryness followed by redissolution into a smaller volume of solvent. A desirable column material is silica.

The column carrying the desired ester compound may then be eluted, for

65

ı is 'n,

уlу

	avamala vaina 1	
	example using a solvent of suitable polarity, e.g. in the case of silica columns, ethyl acetate containing a hydrocarbon such as hexane or toluene.  Finally, the fractions containing the desired at the case of silica columns, ethyl acetate or toluene.	
	Finally, the fractions containing the desired ester compound may be combined and the desired compound obtained by crystallisation. By the foregoing	
5		
		5
	Alternatively, where a carboxylic acid of the investigation	
10		
		10:
	The base may be an organic solvent-soluble base such as sodium 2- ethylhexanoate which will give a salt product. Alternation	10
	treated with an aqueous solution of a water callella lively, the extract may be	
	treated with an aqueous solution of a water-soluble base to form an aqueous solution of the desired salt. In the latter case, this may be further purified by	
15	chromatography, for example, on an anion-exchange resin in the salt, e.g. chloride, form, A suitable resin is AGLY2 (Rio Rod Laborateria in the salt, e.g. chloride,	
	form. A suitable resin is AG1-X2 (Bio-Rad Laboratories, Richmond, California).	15
	The acid of the invention is retained on this room and may be eluted by a salt	
	gradient, the anion of which desirably being the same as that already present on the	
-20	column, and the cation of which being that of which it is desired to prepare a salt of the invention. Thus, for example, the sodium salt of the	
	the invention. Thus, for example, the sodium which it is desired to prepare a salt of with aqueous sodium chloride at gradually into the desired acid can be eluted	20
	with aqueous sodium chloride at gradually increasing concentration, e.g. from 0.1M to 0.25M.	
	Eluted fractions containing the desired company in sale c	
25		
		25
30		
		30
	Where the acid of formula I or a salt thereof has been isolated but an ester thereof is required, the acid or salt may be subjected to esterification. Similarly, where a particular ester has been isolated but an ester	- •
26		
35		2.0
	2	35
	Such deesterification may conveniently be carried out by reductive	
40		
		40
	rhodium. The catalyst may be supported, e.g. on charcoal or kieselguhr. In the case of p-nitrobenzyl esters, cleavage may be effected by reduction of the nitro group (e.g. using a dissolving metal reducing a case).	
45		
45	aqueous tetrahydrofuran or acetone controlled in the pH range 3—6, preferably	45
	4—5.5, by the addition of aqueous HCl; aluminim amalgam in a moist ether e.g.	43
	tetrahydrofuran; or iron and ammonium chloride in an aqueous ether e.g.	
50		
	The alkyl, alkoxyalkyl and aralkyl esters may be prepared by reaction of the acid of formula I with the appropriate diazoalkane or diazo-aralkane, e.g.	50
	The view of the control of the contr	
55		
33		55
		33
60		
	amide solvent, e.g. dimethyl sulphovide dimethyls	60
	hexamethylphosphoramide.	
	Compounds of the invention and methods for their properties and include	
65	"" "O" OC GCSCHOCG III HIE HOHOWING HON-HONITING EVARONICA	
0.5	In the Examples which follow the following steam-sterilised media were used:	65
		0,5

•	8	9	1,585,661	9
as, ethyl			In relation to infrared spectra the symbols s, m and w refer to strong, medium and weak intensity respectively.	
mbined				
regoing			Medium A	
as white	5	5	Medium A Soya bean meal 5 g/l, yeast extract 5 g/l, tryptone 5 g/l, K₂HPO₄ 0.2 g/l, glycerol 10 g/l and tap water to 1 litre.	5
and it is				
acted as		5	Medium B Glycerol 35 g/l, citric acid 1.5 g/l, L. asparagine 6.7 g/l, MgSO <sub>4</sub> . 7H <sub>2</sub> O 0.5 g/l,	
o form a				
	10	Ĭ,	FeSO <sub>4</sub> . 7H <sub>2</sub> O 0.03 g/l, MnSO <sub>4</sub> . 4H <sub>2</sub> O 0.1 g/l and distilled water to 1 litre.	
lium 2-		[ .	FeSO <sub>4</sub> . /H <sub>2</sub> O 0.03 g/l, MnSO <sub>4</sub> . 4H <sub>2</sub> O 0.1 g/l and comme	
may be	·	_	14 E	10
aqueous		] 10	Medium C Sucrose 20 g/l, distillers solubles 15 g/l, yeast extract 0.2 g/l, K <sub>2</sub> HPO <sub>4</sub> 0.2 g/l,	• -
ified by		Į	tryptone 5 g/l, glycerol 10 g/l and tap water to 1 litre.	
hloride,	15	[	tryptone 5 gr, giveror to gr and tap water to 1 more	
ifornia).		}	Madium D	
y a salt			Medium D Soya meal (unmilled) 30 g/l, ferric sulphate 0.1 g/l, KH <sub>2</sub> PO <sub>4</sub> 0.1 g/l, soluble	
it on the		}		15
a salt of	20	15	The silica used in column chromatography procedures is Woelm silica (activity	
e eluted g. from	20	}	grade III).	
.g. HOIII				
ner with		}	Example 1	
nich will			2-Hydroxymethyl-clavam	20
passage	-25	20	a) Fermentation of Streptomyces clavuligerus strain	20
Isorbent	23	200	a) Fermentation of Streptomyces clavaligerus strain NCIB 11261 was maintained on malt agar slopes Strepomyces clavaligerus strain NCIB 11261 was maintained on malt agar slopes Strepomyces clavaligerus strain NCIB 11261 was maintained on malt agar slopes Strepomyces clavaligerus strain NCIB 1261 was maintained on malt agar slopes Strepomyces clavaligerus strain NCIB 1261 was maintained on malt agar slopes Strepomyces clavaligerus strain NCIB 1261 was maintained on malt agar slopes Strepomyces clavaligerus strain NCIB 1261 was maintained on malt agar slopes Strepomyces clavaligerus strain NCIB 1261 was maintained on malt agar slopes Strepomyces clavaligerus strain NCIB 1261 was maintained on malt agar slopes Strepomyces clavaligerus strain NCIB 1261 was maintained on malt agar slopes Strepomyces clavaligerus strain NCIB 1261 was maintained on page 1660 was malt agar slopes strain NCIB 1261 was maintained on page 1660 was malt agar slopes strain NCIB 1261 was malt agar s	
miscible		1	(malt extract 24 g/l; yeast extract 5 g/l; agai 15 g/l; adjusted to pri 1107 g/l	
icetone.		ţ	weeks at 28°C.	
ined by		}	weeks at 28°C.  The slopes were developed for shake flask fermentation with 1/3 of a slope being used to inoculate 50 ml of Medium A (pH adjusted to 6.5) in a 250 ml flask.	25
	30	25	being used to inoculate 50 mil of Medalin A 200 row/min on a rotary shaker with a 2"	
an ester			This was incubated at 28°C for 42 h at 220 rev/min on a rotary shaker with a 2" throw. 2 ml of the inoculum was used to inoculate 50 ml of Medium B containing throw. 2 ml of the inoculum was used to inoculate 50 ml of Medium B containing throw. 2 ml of the inoculum was used to inoculate 50 ml of Medium B containing throw.	
milarly,		}		
e initial			NaCl (0.1 g/l) and uracli (0.01 g/l) (pH adjusted to 7.0 with a 5 cm throw unbaffled flask. This was kept at 28°C for 72 h on a rotary shaker with a 5 cm throw	
ase, the	3.5		at 220 rev/min.	30
i as an	35	30	•	
ductive		1	b) Isolation of 2-hydroxymethyl-clavam  Broth (51) prepared in a) above was centrifuged and the supernatant applied to	
ige, e.g.		}		
rs. Such			a column containing XAD-2 resin (bed lit. 100 cm. dain state) acetate (3×1 litre). saturated with ammonium sulphate and extracted with ethyl acetate (3×1 litre).	
lyis, e.g.	40	1 25	The extracts were combined, dried over sodium sulphate and evaporated to	35
dium or		35	dryness under reduced pressure.	
the case		ļ		
o group		1		
r zinc in eferably	45	i	evaporated under reduced pressure. The resultant some eluted with toluene-ethyl column containing dry silica (30×1 cm) and the column eluted with toluene-ethyl column containing dry silica (30×1 cm) and the reactions 11—15 were combined and	40
her e.g.	43	40	column containing dry silica (30x1 cm) and the column containing dry silica (30x1 cm) and the column containing dry silica (30x1 cm) and the column c	
such as		1	acetate (1:1), 20 ml fractions being confected. Tactions as a yellow oil (40 mg), evaporated under reduced pressure to give the alcohol as a yellow oil (40 mg).	
:duction		ļ	The infrared spectrum of a bromoform solution of the alcohol showed peaks (cm <sup>-1</sup> ) at 3660 w, 3560 m, 2930 n, 1770 s, 1600 w, 1455 w, 1392 m, 1332 s, 1233 m,	
n of the	50	Ï		45
ne, e.g.		45		
ntly be		J	The infrared and nmr spectra are shown in Figs. 2 and 1 respectively.	
int, e.g.		Ĩ	The initiated and initi openio are as	
ethane.		<b>↓</b> .	Example 2	
reactive	55	J	2-Hydroxy-methyl-clayam	
mide or		1 60	p it (21) proposed as in Example 1(2) above was centrifuged and the	50
r, with a ich as a		. 50	and the second second to a column containing XAD-4 lesin (bed neight so they	
such as	•		11 WALEL TE am I The feels was washed will water, then close with me	
oxide or	60	Ϋ́	water (1:1 by vol.). The cluate was evaporated under reduced pressure to an	
de or		<b></b>	1 4	. 55
		55	mi 1 i.m. / 5/1 mill tilde evitacien with 4x 1/1 iii Eliivi decidie und	,,,
solation	•	7	extracts were combined, dried over 142,504 and evaporated and	
		J.	pressure to a residue.	
re used:	65	1	The residue was dissolved in a little ethyl acetate, silica was added and the	
		. <del> </del> .	·	

	mixture evaporated under reduced	10-
	mixture evaporated under reduced pressure. The resultant dry silica was applied to a column containing dry silica (Woelm, act. grade III, 20×1 cm). Elution was with toluene-ethyl acetate (1:1) and 20 ml fractions were collected. Fractions 10—12 were combined and evaporated to give 13 mg of a pale-yellow oil.	
5	The product had at a con-	
	The product had the following properties  TLC. Portions of the product were subjected to TLC on silica with the solvents given below. Assays were by overlay with Saccharomyces carlsbergensis.	5:
	Solvent	
10	Methanol EK6061 (0 Imm thick plactic backet) Rf	
10	EK0001	
	Merck 60 (plastic backed) 0.62 Chloroform EK6061 0.40	10
	Merck 60 0.26	
15	Toluene EK6061 0.04	
	Merck 60° Toluene-ethyl acetate EK6061 0	1.5
	(1:1) Merck 60 0.41	
	Toluene-methanol EK6061 0.18 (9:1) Merck 60 0.40	
•	(9:1) Merck 60 0.40 0.15	
20	(E. Merck is a registered Trade Mark)	
		20
	Addition of 2 days Son Son UV Spectrum	
	absorption with 3 270 -m NaOH to a methanolic solution resulted in a strong	
25	I he intrared and nmr spectra ware circulations and the intrared and nmr spectra ware circulations.	
23	1. Spectra were similar to those of the product of Example	<u> </u>
	Example 3	25
	a) Inoculum development	
30	Streptomyces clavuliganus NCID 11200	
30	The contents of one ampoule was suspended in sterile distilled water and then added to Medium C adjusted to pH 6.5 with hydrochloric acid.	
	added to Medium C adjusted to pH 6.5 with hydrochloric acid in a 250 ml shake	30
,	26°C for 72 h. The start of the	
35	A portion (2 ml) of the 72 h inoculum was used to inoculate each of 4×2 litre	
	incubated on a rotary shaker at 220 revision in the florence flasks were	35
	The contents of three of the florence flasks (3×150 ml 3.75 v/v) were used to inoculate 3×5 litre fermenters each containing 4 litera of Mal.	55
	6.5 with NaOH/HCI) advantage intestit Medium D (adjusted to nH	
40	These fermenters were agitated with 2027 dis	·
	rev/min at 28°C for 20 h with an air flow of 3 litres/min.	40
	INOCHINITO I / 3 litron 50/ m/s /	
	150 litres of the above soya meal medium in a 230 litre stainless steel fermenter.  The vessel was agitated with a six bladed 8" impeller at 250 and a 250 a	
45	The vessel was agitated with a six bladed 8" impeller at 350 rev/min and aerated at 420 litre/min for 20 h at 28°C.	
•	b) Fermenation	45
	Broth from the 230 litre formands (60 th	
•	Broth from the 230 litre fermenter (50 litres, 10% v/v) was used to inoculate 430 litres of Medium D in a 700 litre stainless steel fermenter. The vessel was 250 rev/min and accounted to the control of	
50	agitated with a six bladed 10" impeller at 350 rev/min and aerated at 280 litre/min.  The fermentation was carried out at 28°C for 92 h, and was residued.	
	The fermentation was carried out at 28°C for 92 h, and was maintained at pH 6.5.	50
	c) Extraction of fermentation broth	50
	marvest broth (530 1 mt cas a	
	sulphuric acid, filter aid (15 kg) added, and the mixture clarified on a rotary drum	
55	CAL, 105 l) in columns. The carbon was adsorbed onto carbon (Pittsburgh	
	were eluted with aqueous acetone (60%) with the water (70 I) and the columns	55
	The combined eluates (148 1) were applied to a column of IRA68 resin	
	· · · · · · · · · · · · · · · · · · ·	

	10:2	11		1,585,661		11
pplied to			(ablacida avala 12.4.1) the officer	1 11 11		11
was with	- 1	ł	(chloride cycle, 12.4 1), the effluent water (20 1).	being collected. The	column was washed with	
s 10—12	100		water (20 1).			
solvents	Service Services	5	d) Isolation of 2-hydroxymethyl-cla The combined effluent and was in a pot still to 50 1, and the pH adjus solution. A portion (25 1) of this sulphate and stirred with ethyl aceta	shings (148 1) from st sted to 7.0 with 40% ac concentrate was sai	queous sodium hydroxide turated with ammonium	5
Rf 0.79 0.62	10 <sup>温</sup>	10	concentrated under reduced pressur A portion of this oil (13.6 g) was	to with annydrous noted to yield the title control of the control	nagnesium sulphate and ompound (46 g) as an oil.	10
0.40 0.26 0.04 0	10		and washings were applied to a colu LH20 packed in ethyl acetate. The fractions being collected. Fractions	mn (bed ht. 53 cm, di column was eluted v	cetate. Combined filtrate iam. 5.2 cm) of Sephadex with ethyl acetate, 50 ml	10
0 0.41 0.18 0.40	15	15	A portion of this oil (960 mg) was acetate (1:1; 2 ml) and applying to a silica powder. The column was dev	From 2/3, M/10 NaoH Sfurther purified by d dry column (bed ht.	issolving in toluene-ethyl 23 cm, diam. 2.1 cm) of	15
0.15	20	20	(1:1), the eluate being collected in collected the eluting solvent was characteristic combined and evaporated to give the The i.r. and n.m.r. spectra of the obtained for the product of Example solution of the compound in M/10 N	e title compound as a perioduct were substa	e. Fractions 45—73 were pale yellow oil (220 mg). Initially the same as those	20
strong	}	25	solution of the compound in M/10 N sulphoxide -166°.	aon was 841 at $\lambda_{\text{max}}$	259 nm. $[\alpha]_0^{23}$ in dimethyl	25
ımple	1		·	Example 4		
····pic	25		a) Fermentation of Streptomyces cla Streptomyces clavuligerus strain	oxymethyl-clavam vuligerus strain NCIB 11261 was m	naintained on malt agar	
ules. then hake	30	30	for 2 weeks at 28°C. Sterile water (8 made. 2ml portions of this suspensio baffled shake flasks containing 50 m (pH adjusted to 6.5 with HCl).	ct 5g/l;, agar 15g/l; acml) was added to eacn n were used to inocu l of Medium A conta	djusted to pH 7.8) grown h slope and a suspension late each of four 250 ml aining uracil (100 µg/ml)	30
litre vere h.	35	35	The flasks were incubated on throw, at 26°C for 24 h. The shake floof this inoculum used to inoculate 3 above medium and incubated for 48 v/v) portions of this bulked 48 h	2×250 ml baffled sha h under the previous	ake flasks containing the sconditions. 120 ml (3%)	35
d to pH 750	40	40	additionally containing varying level The uracil concentration and the	s of uracil. ne aeration rate in e	usted to 7.0 with NaOH)	<b>40</b>
ate er. at		45	5 litre Fermenter Uracil No. 1 2	concentration μg/ml 100 100	Aeration rate litre/min. 1.5	45
ıte	45	-· 50	3 4 5	100 100 50 35	3 6 6 6	
as	Ţ	٠	6	25	6	50
n. 5. 5	50 J.		The 5 litre fermenters were agin bladed impellers. The fermentations	were maintained at	with 2×3½" diameter 4 28°C for 94 h.	
th m th	5	55	b) Isolation of 2-formyloxymethyl-cla Bulked broth from fermenters supernatant applied to a column conta cm). The resin was washed with water 500 ml). The eluate was evaporated adjusted to 6.8 with sodium bydrovice.	l—6 above (18 1) waining XAD-4 resin (b (4 1) and eluted with	ped ht. 130 cm; diam. 2.8 acetone-water (4:1) (c.	55
n		60	adjusted to 6.8 with sodium hydroxi sulphate and extracted with ethyl aced dried (sodium sulphate), evaporated	ate (4×100 ml) The	urated with ammonium	60

12	1,085,086,1	12
5	dissolved in a little ethyl acetate. Silica was added and the mixture evaporated under reduced pressure to give a solid which was added to the top of a column containing dry Woelm silica (act. grade III; bed ht. 50 cm; diam. 2.9 cm). The column was eluted with toluene-ethyl acetate (1:1), 15 ml fractions being taken. Fractions 4—11 were combined, evaporated under reduced pressure and the residue subjected to chromatography on silica (bed ht. 30 cm; diam. 2.1 cm), with hexane-ethyl acetate (2:1) as eluant, 10 ml fractions being collected. Fractions 17—23 were combined and evaporated under reduced pressure to give the formyloxy	
10	solution of the sample is shown in Fig. 3 of the accompanying drawings and has peaks (cm <sup>-1</sup> ) at 2915 m, 1784 s, 1730 s, 1462 w, 1406 w, 1392 w, 1346 m, 1282 m, 1066 s, 1034 s, 990 m, 940 w, 926 w and 863 w. A 100 MHz proton nmr spectrum of a solution in deuterochloroform had $\tau$ values as shown in Table 2	10
15	Broth (50 µl) from 5 litre fermenter No: 6 was applied to Merck silica 60 F254 plates, glass-backed, and the plates developed with ethyl acetate, air-dried and overlayed with nutrient agar containing Saccharomyces carlsbergensis 1738. The formyloxy compound was found to have an R, value of 0.89. The formyloxy compound was also detected by quenching of fluorescence under u.v. light (254 nm) after treatment of the developed TLC plate with ammonia vapour.	10 10 15 15 15 15 15 15 15 15 15 15 15 15 15
20	Example 5	20
	2-Formyloxymethyl-clavam Filtered broth (25 µl) obtained as in Example 3 was applied to Merck silica 60 F254 plates, glass-backed, which were developed with ethyl acetate and air-dried. The plates were then saturated with ammonia vapour and the chromophore	
25	produced on reaction with ammonia detected under u.v. light (254 nm). The formyloxy compound was found to have an R <sub>t</sub> value of 0.85.	25
	Example 6 2-Formyloxymethyl-clavam	
	a) Inoculum development	
30	Streptomyces clavuligerus strain NCIB 11261 was maintained on malt agar slopes (malt extract 2.4%; yeast extract 0.5%; agar 1.5% w/v; pH 7.8) grown for 2 weeks at 28°C. Tween 80 solution (8 ml) (Tween is a registered Trade Mark) was added to the slope and a suspension made. 2 ml portions of this suspension were used to inoculate each of four 250 ml baffled shake flasks containing 50 ml of	30
35	Medium A containing uracil (100 μg/ml) (pH adjusted to 6.8—7.1).  The flasks were incubated on a rotary shaker at 220 rev./min with a 2" throw, at 28°C for 48h. 2 ml portions of this inoculum were used to inoculate identical shake flasks containing the above medium and incubated for 24 h. under the previous conditions. The shake flask contents were bulked and 120 ml portions transferred	35
40	to 250 ml aspirators. These 120 ml (3% v/v) portions were then used to inoculate 5 litre fermenters each containing 4L of Medium B containing NaCl (0.1 g/l), antifoam (0.05% v/v) and uracil (35 μg/ml), (pH adjusted to 7.0 with KOH).  The fermenters were agitated at 250 rev/min with 2×3½" diameter 4 bladed impellers. The fermentations were maintained at 28°C for 24 h. at an aeration rate of 6 l/min.	40
	7.51 (5% v/v) of this 24 h culture were used to inoculate a 230 litre fermenter containing 150 1 of the above medium. The fermentation was maintained at 28°C for 93 h. at an aeration rate of 5 cu. ft/min and an agitation rate of 250 rev/min. Further antifoam was added as required throughout the fermentation.	45
50	b) Isolation of 2-formyloxymethyl-clavam (i) Harvest broth (133 l) was adjusted to pH 7.0 with sodium hydroxide and clarified by centrifugation. To the clear supernatant (114 l) 1/3 volume butanol was added, stirred for 30 mins and the phases separated by centrifugation. The aqueous	50
55	butanol extracts (76 1) washed with distilled water (40 1).  Distilled water (40 1) was added to the butanol extracts and the azeotrope concentrated to 11 by means of a pot still and evaporation under reduced pressure.  The concentrate was saturated with ammonium sulphate and extracted with 4.	55 ·
60	volume ethyl acetate. The bulked ethyl acetate extracts (1.95 1) were dried with magnesium sulphate, evaporated to 500 ml, dry silica added (Sorbsil M60, 50 g) and evaporated to dryness.	60 .

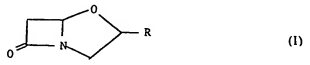
	12	13	1,585,661	13
orated olumn ). The .aken.			The solid residue was added to the top of a column containing dry silica (Sorbsil M60, 142×3.8 cm) (Serbsil is a registered Trade Mark) and the silica was eluted with toluene/ethyl acetate (1:1), 25 ml fractions being collected. Fractions 38—73 inclusive were bulked.	
id the with 17— yloxy oform	<b>5</b> 	5	(ii) Harvest broth (133 1) prepared as in (a) above was clarified by centrifugation and the supernatent (123 1) evaporated to 221 on a pot still. The concentrate was adjusted to pH 7.0 with sodium hydroxide, filtered, saturated with ammonium sulphate and extracted with 2x1 volume ethyl acetate. The bulked ethyl acetate extracts (21.5 1) were washed with saturated ammonium sulphate	5
d has 1066 of a F254	10 .	10	solution (5 1), dried with magnesium sulphate and evaporated to 250 ml under reduced pressure. Sorbsil silica (50 g) was added, the mixture evaporated to dryness and the solid residue added to the top of a column containing dry silica (Sorbsil M60, 138×3.8 cm). The silica was eluted with toluene/ethyl acetate (1:1), 25 ml fractions being taken. Fractions 42 to 60 were bulked.	10
and The loxy (254	15	15	(iii) The bulked eluates from (i) and (ii) were combined, evaporated to 15 ml and applied to a column containing LH20 Sephadex (60x60 cm) in ethyl acetate. The Sephadex was eluted with ethyl acetate, 25 ml fractions being collected. Fractions 38—46 were bulked and evaporated to dryness to yield title compound (700 mg) having characteristics similar to those described in Example 4.	· 15
	20	20	Example 7	20
a 60 ied. :ore		·	Methyl clavam-3-carboxylate  a) Extraction of fermentation broth Harvest broth (520 1, pH 6.7) obtained as in Example 3 was adjusted to pH 5.4	
The	25	25	with sulphuric acid, filter aid (15 Kg) added, and the mixture clarified on a rotary drum filter with a precoat. The filtrate was adsorbed on to carbon (Pittsburgh CAL, 105 1) in columns, the carbon was washed with water (70 1) and the columns were eluted with aqueous acetone (60% v/v, 135 1).  The combined eluates (143 1) were applied to a column of IRA 68 resin	25
gar r 2 vas	30	30	(chloride cycle, 12.5 1). The column was washed with water (20 1) and eluted with 5% (w/v) aqueous lithium chloride solution (16 1), collecting the eluate in fractions of 1 1. Fractions 6—15 were combined and cold propan-2-ol (5 vol.) was added with stirring, followed by filter aid (500 g) and the mixture filtered. The cake was washed with water/propan-2-ol (1:5, 2—4 1) and the combined filtrates concentrated under	30
of at	35	35	reduced pressure to 1/6 volume of combined eluates. The resulting concentrate was stored at 4°C for 16 h, during which time a solid formed. The solid was removed by filtration and washed with 30% aqueous lithium chloride solution (3×50 ml).	35
ke us ≥d · 5 !), :d :e	40	40	b) Isolation of methyl clavam-3-carboxylate  Mother liquor and washings from step (a) (2 1, pH 5.8) were saturated with about 800 g of ammonium sulphate, the pH adjusted to 3.0 with 11.4M hydrochloric acid and extracted with ethyl acetate (2×500 ml). The ethyl acetate extracts containing the β-lactam acid were combined, dried over sodium sulphate and treated with excess diazomethane prepared by the method of Mlejnek (J. Chromatog. 1972, 70, 59).	40
er	45	45	The ethyl acetate extract was evaporated under reduced pressure, the residue dissolved in a little ethyl acetate, silica added and the mixture evaporated under reduced pressure. The resultant solid was added to the top of a column containing dry silica (bed ht. 50 cm; diam. 2.9 cm) and the column eluted with hexane-ethyl acetate (1:1 by vol.), 20 ml fractions being collected. Fractions 8—14 were	45
d s	50	50	combined and evaporated under reduced pressure to yield a yellow oil (810 mg), which was dissolved in a little ethyl acetate, silica added and the mixture evaporated under reduced pressure. The resultant solid was added to the top of a column containing dry silica (bed ht. 40 cm; diam. 2.5 cm). The column was eluted with hexane-ethyl acetate (3:2 by vol.), 10 ml fractions being collected. Fractions 22—28 were combined and evaporated under reduced pressure to yield the methyl	50
i :	55	55	ester as a pale yellow oil (200 mg). Found: C, 49.1 49.3; H, 5.3, 5.3; N, 7.7, 8.1%; S, nil. C,H <sub>9</sub> NO <sub>4</sub> requires C, 49.1: H, 5.3; N, 8.2%. The infrared spectrum of a bromoform solution of the sample showed absorption peaks at 2950, 1785, 1754 and 1215 cm <sup>-1</sup> . A 100 MHz proton n.m.r. spectrum of the ester in deuterochloroform solution had $\tau$ values of 4.52 (d, 2.5) (1H); 5.16 (dd, 4.5, 7.5) (1H): 5.88 (dd, 7.5, 11.5)	55
	60	60	and 6.90 (dd, 4.5 11.5) (2H); 6.24 (s) (3H); 6.69 (dd, 2.5, 16) and 7.14 (d, 16) (2H) and this is shown in Fig. 6.	60

And the second s

.....

#### WHAT WE CLAIM IS:

#### 1. A compound of the formula (I)



wherein R represents a hydroxymethyl group, said compound in deuteroacetone exhibiting the 100 MHz proton n.m.r.  $\tau$  values shown in Table I herein, and the 5 corresponding compounds in which R is a formyloxymethyl group or a carboxyl or esterified carboxyl group or, in the case where R represents a carboxyl group, salts thereof. 2. The compound as claimed in claim 1 wherein R represents a hydroxymethyl 10 group. 10 The compound as claimed in claim I wherein R represents a formyloxymethyl group. 4. The compound of claim 2 which has a negative molecular rotation [M]2 in dimethyl sulphoxide of -166° or the formic acid ester thereof. 5. The compound as claimed in claim 3, a deuterochloroform solution of which 15 15 has  $\tau$  values in the 100 MHz proton nmr spectrum as shown in Table 2 herein. 6. A compound as claimed in claim 1 wherein R represents a carboxyl or esterified carboxyl group, or when R is a carboxyl group, salts thereof. 7. The acid as claimed in claim 6, the diphenylmethyl ester of which in a deuterochloroform solution has  $\tau$  values in the 100 MHz proton nmr spectrum as 20 20 shown in Table 3 herein. 8. The acid as claimed in claim 6 or claim 7, the diphenylethyl ester of which has a negative molecular rotation [M]23 in dimethyl sulphoxide of -352°. 9. An ester of the acid as claimed in any of claims 6 to 8, wherein R represents an esterified carboxyl group -COOR' wherein R' represents a straight or 25 branched substituted or unsubstituted alkyl or alkenyl group having from 1-8 25 carbon atoms; an aralkyl group having up to 20 carbon atoms; an aryl group having up to 12 carbon atoms; or a cycloalkyl group containing up to 12 carbon atoms optionally containing one or more heteroatoms in the ring system, and 30 unsaturation optionally being present when a heteroatom is present. 10. A compound as claimed in claim 9 wherein R1 represents a C1-4 alkyl 30 group. 11. A compound as claimed in claim 9 wherein R' is a methyl group. 12. A compound as claimed in claim 9 in which R1 is an arylmethyl group. 35 13. An alkali metal, alkaline earth metal, ammonium or organic base salt of the 35 carboxylic acid claimed in claim 6, claim 7 or claim 8. 14. The sodium, potassium, lithium and magnesium salts of the carboxylic acid claimed in claim 6, claim 7 or claim 8. 15. A pharmaceutical composition comprising an antifungal compound as claimed in any of claims 1-14 in which R is a hydroxymethyl, formyloxymethyl or 40 40 esterified carboxyl group and one or more pharmaceutical carriers or excipients. 16. A composition as claimed in claim 15 in a form suitable for oral, topical, rectal, intravaginal or parenteral administration. 17. A composition as claimed in claim 15 or claim 16 in a form suitable for 45 topical administration. 18. A composition as claimed in any of claims 15-17 containing the active 45 compound at a concentration of from 0.1 to 95% by weight. 19. A composition for horticultural or agricultural use which comprises one or more antifungal compounds as claimed in any of claims 1-14 in which R is a hydroxymethyl, formyloxymethyl or esterified carboxyl group in association with a 50 carrier or diluent. 20. A composition as claimed in claim 19 containing the active material at a concentration of from 0.01 to 40% by weight. 21. A composition as claimed in claim 19 or claim 20 in the form of a dust, granulate, powder, pellet, spray, smoke or mist.

22. A process for the preparation of a compound of formula (I) as claimed in 55 claim I which comprises isolating the compound of formula (I) wherein R is a hydroxymethyl, formyloxymethyl or carboxyl group from a fermentation broth of a

strain of Streptomyces clavuligerus or where an ester of formula (1) is required,

therefrom of fractions containing said acid.

resin as defined in claim 28.

55

65

43. A process as claimed in claim 42 in which the anion exchange resin is a

44. A process as claimed in claim 42 or claim 43 wherein removal of clavulanic

		18
	acid is effected by reaction of the material containing the acid of formula (1) or a salt thereof with a lithium salt to form the lithium salt of clavulanic acid and separating said salt by fractional crystallisation or precipitation.	5
· 5		Ñ
	acidic compound of formula I is adsorbed on a column and is eluted with an aqueous solution of a lithium salt or is eluted with an addition of a lithium salt to the eluted	5
	addition of a lithium salt to the eluate.	₹,
	46. A process as claimed in claim 45 mb :	-
10	at high concentration is incorporated into the solution of the lithium salt used for the elution or into the agueous eluque to essist soid for the lithium salt used for	
	the elution or into the aqueous eluate to assist aid fractional crystallisation or precipitation in order to provide 70. 07% best land fractional crystallisation or	10
	47. A process as claimed in claim 46 by volume of said solvent.	
		• •
15	tetrahydrofuran, dimethylformamide or dimethylsulphoxide.	
13	70. A DIOCESS AS CIAIMED IN ANY At claime 43 to 47 to 11 to 1	16
	of said carboxyl compound onto an anion exchange resin it is adsorbed onto	15
	adsorbent carbon and eluted therefrom with an aqueous water-miscible solvent.  49. A process as claimed in any of claims 44—48 wherein the compound of formula (I) wherein R represents a carbon of the compound of the compou	
20		
20 -		20
		20
	50. A process as claimed in any of claims 41 to 49 in which compound of formula (I) in which R is an esterified carboxyl group is formed and the compound in which R is a carboxyl group is obtained therefore is formed and the compound	•
25		0.5
		25
	52. A process as claimed in any of claims 41 to 49 in which the acid of formula  (I) in which R is a carboxyl group is esterified without isolation of said acid.	
30		•
		30
	54. A process as claimed in any of claims 41—52 wherein a salt of a compound of formula (I) in which R is carboxyl is esterified by reaction with reactive derivative of an alcohol	
	derivative of an alcohol.	
35	55. A process as claimed in claim 52 in which esterification is effected on impure material of formula (I) where P is combanily and the state of the	26
		35
	purified by chromatography on silica.  56. A process as claimed in claim 22 substantial.	
	56. A process as claimed in claim 22 substantially as hereinbefore described.  57. A process as claimed in claim 22 substantially as hereinbefore described with reference to the Examples	
40		40
	58. A compound of formula (I) as shown in claim I wherein R is a	40
	hydroxymethyl, formyloxymethyl, carboxyl group or esterified carboxyl group or, when R is a carboxyl group a salt thereof in h	
	claimed in any of claims 22-57	
45	59. The compound of claim 2 substantially forms	
	60. The compound of claim 3 substantially free of any isomeric material.  61. A compound of claim 6 substantially free of any isomeric material.	45
	61. A compound of claim 6 substantially free of any isomeric material. 62. The compound of claim 2 whenever produced by the claim 2 whenever produced by the claim 2 whenever produced by the claim 2 whe	
	Streptomyces clavilinerus	
50	63. The compound of claim 3 whenever produced by the c	
		50
	64. A compound of claim 6 wherein the compound in which R represents a	
÷	together with esters or salts thereof	
55	65. The compound of claim 2 substantially 6.	
	derived materials.	55

Alidabah dikipatin ini birantar baran di bada birin barapa da

I) or a id and le said ith an red by olvent ed for on or 10 ganic oxan,

ption 15 onto vent. nd of other lvent 20

id of ound

ed is 25

d of ith a 30 i

on 35

red. ped 40

; a 40 or, as

45

of 50

· a · s,

n- 55

66. The compound of claim 3 substantially free from other fermentation-derived material.

67. A compound of claim 6 wherein R represents a carboxyl group substantially free from other fermentation-derived material.

For the Applicant, FRANK B. DEHN & CO., Chartered Patent Agents, Imperial House, 15—19 Kingsway, London, W.C.2.

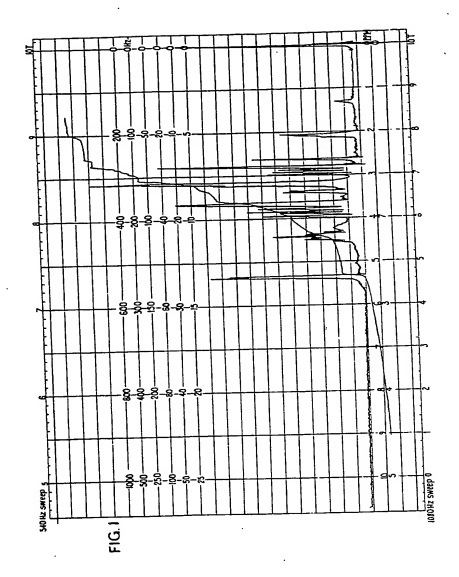
Printed for Her Majesty's Stationery Office, by the Courier Press, Learnington Spa, 1981 Published by The Patent Office, 25 Southampton Buildings, London, WC2A 1AY, from which copies may be obtained.

COMPLETE SPECIFICATION

6 SHEETS

This drawing is a reproduction of the Original on a reduced scale

Sheet 1

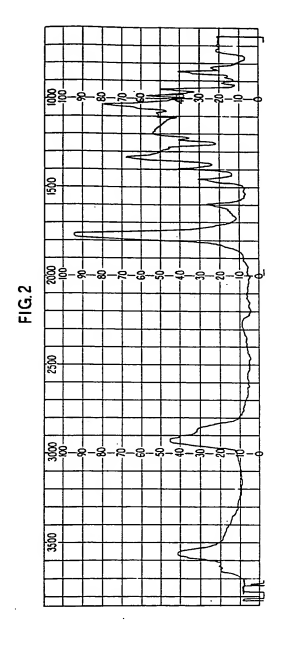


COMPLETE SPECIFICATION

6 SHEETS

This drawing is a reproduction of the Original on a reduced scale.

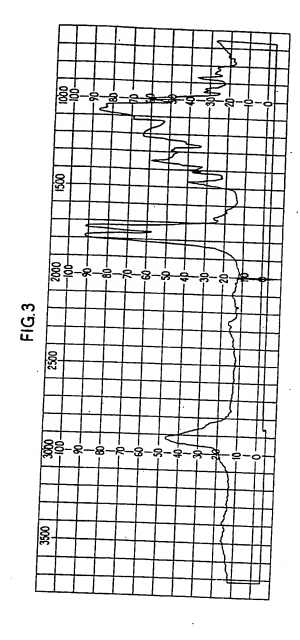
Sheet 2



COMPLETE SPECIFICATION

6 SHEETS

This drawing is a reproduction of the Original on a reduced scale
Sheet 3



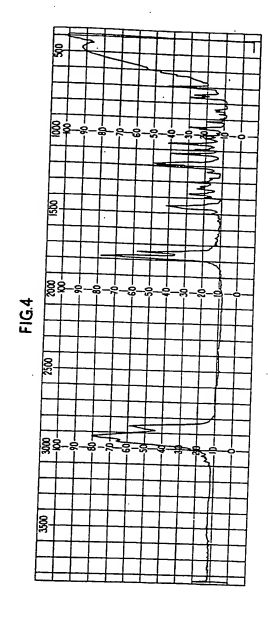
on the front is near technique, with the

COMPLETE SPECIFICATION

6 SHEETS

This drawing is a reproduction of the Original on a reduced scale

Sheet 4



THE RESIDENCE OF THE PROPERTY OF THE PROPERTY

COMPLETE SPECIFICATION

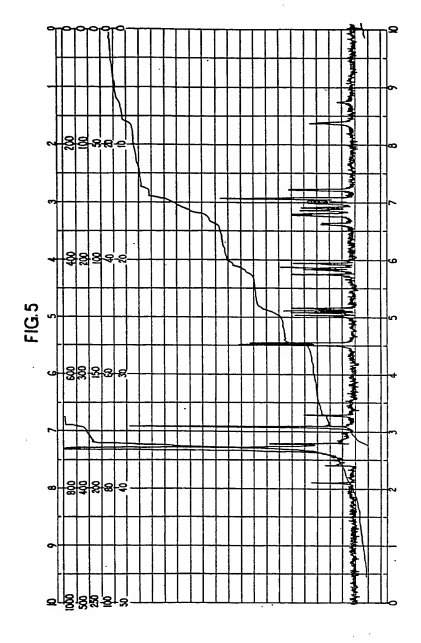
6 SHEETS

This drawing is a reproduction of the Original on a reduced scale Sheet 5

Telegram is a second of the se

The substitution of the state o





COMPLETE SPECIFICATION

6 SHEETS

This drawing is a reproduction of the Original on a reduced scale

Sheet 6

